

REPORT DOCUMENTATION PAGE

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FILE

MEMORANDUM FOR PRS (Contractor Publication)

19 May 2003

FROM: PROI (STINFO)

SUBJECT: Authorization for Release of Technical Information, Control Number: AFRL-PR-ED-VG-2003-138

5364 Boatz, Jerry (PRSP), "New Materials Design"

DoD High Performance Computing Users Group Conference
(Bellevue, WA, 9-13 Jun 2003) (Deadline = 9 Jun 2003)

(Statement A)

New Materials Design

DoD UGC, 9-13 June 2003

Bellevue, WA

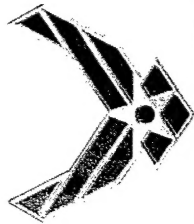


Jerry Boatz

Senior Research Chemist

Propulsion Directorate

Air Force Research Laboratory



NEW MATERIALS DESIGN



THE TEAM....

Prof. Mark S. Gordon

Prof. Gregory Voth

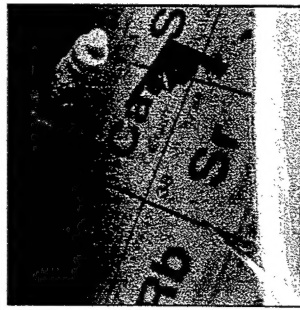


Prof. Sharon Hammes-Schiffer

PENNSSTATE

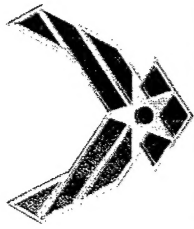


Dr. Ruth Pachter, AFRL/MLPJ



Dr. Jerry Boatz, AFRL/PRSP





OUTLINE



1. Project Overview

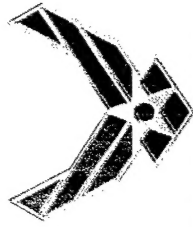
- High energy density materials
- Polyhedral oligomeric silsesquioxanes (POSS)
- Non-linear optical materials

2. Theoretical Methods and benchmarks

- Ab initio electronic structure theory
- Nuclear-electronic orbital approach

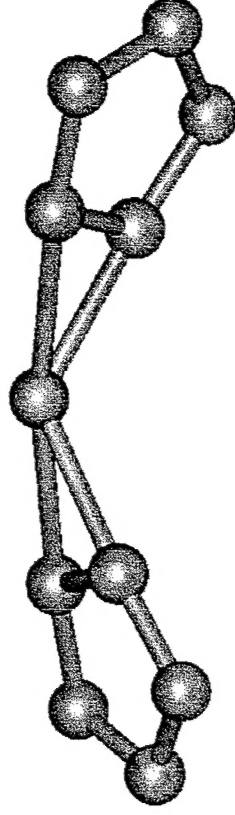
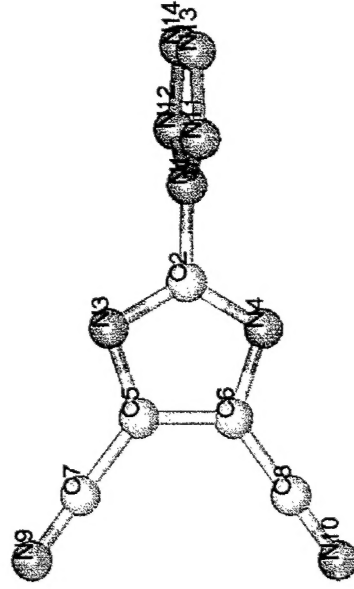
3. Results

4. Summary

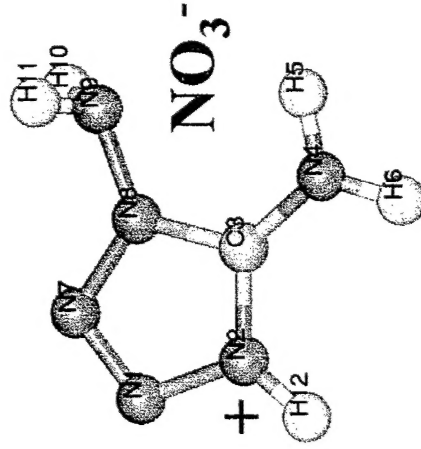


PROJECT OVERVIEW - HEDM

High Energy Density Matter -- next generation rocket propellants



High-nitrogen/polynitrogen compounds

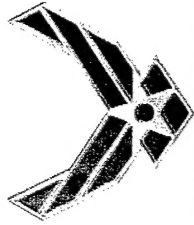


NO_3^-

Energetic Ionic Liquids

Specific Impulse

$$I_{sp} \propto \sqrt{\Delta H / m}$$



PROJECT OVERVIEW - HEDM



Technical issues being addressed using CCM

1. High-nitrogen/polynitrogen compounds

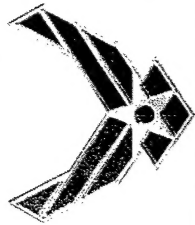
Objective: identify, characterize, and synthesize stable compounds with high heats of formation, high densities

- geometries, energy content, stabilities, reaction pathways

2. Energetic ionic liquids

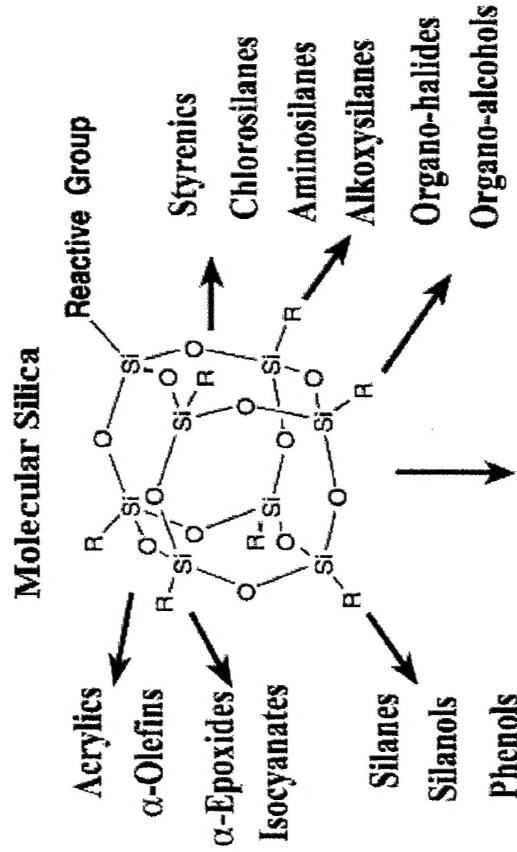
Objective: identify, characterize, and synthesize thermally stable energetic ILS with low melting points, high densities and low viscosities.

- geometries, electronic structures, stabilities, energy content, interaction potentials



PROJECT OVERVIEW - POSS

Polyhedral oligomeric silsesquioxanes -- next generation plastics



As Additives

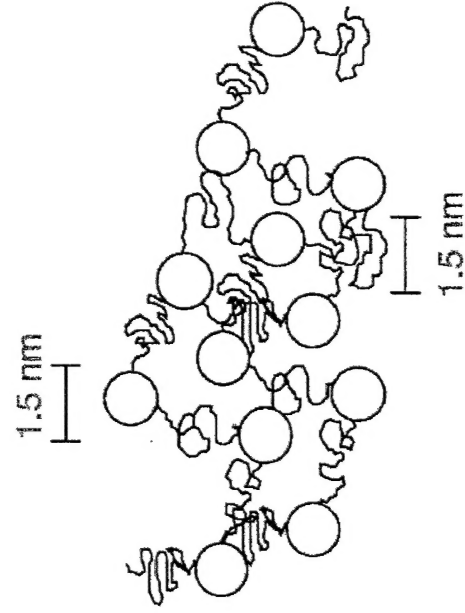
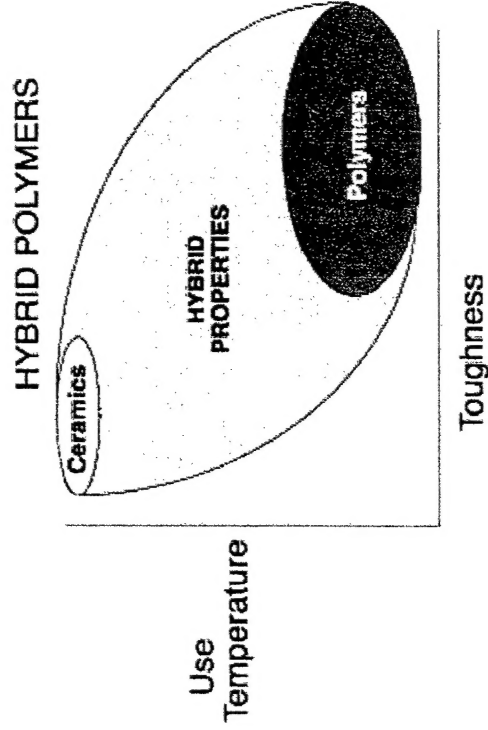
Heat/abrasion resistant paints and coatings
 Mechanical property/viscosity/thermal modifiers
 Crosslinking agents
 Fire retardants

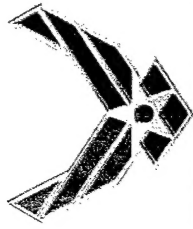
As Plastics

Medical materials
 Space resistant resins
 Packaging/coatings
 Electronic materials
 Optical Plastics

As Preceramics

Ablative materials (nozzles, insulations etc.)
 Claddings/electronics coatings
 Precursors to glassy or ceramic matrices





PROJECT OVERVIEW - POSS



Technical issues being addressed using CCM

1. Mechanisms of formation

Objective: rational design and synthesis of POSS

- role of solvents, acid/base catalysis, substituent effects on mechanism of formation

2. Potential applications as molecular “sieves”

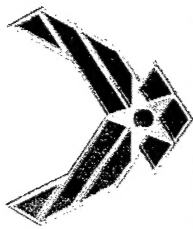
Objective: determine if POSS cages can be used to separate small molecules

- determine barriers to encapsulation of N_2 and O_2

3. Ti-POSS as new catalysts

Objective: determine if Ti-POSS and Ti-siloxane compounds are effective catalysts

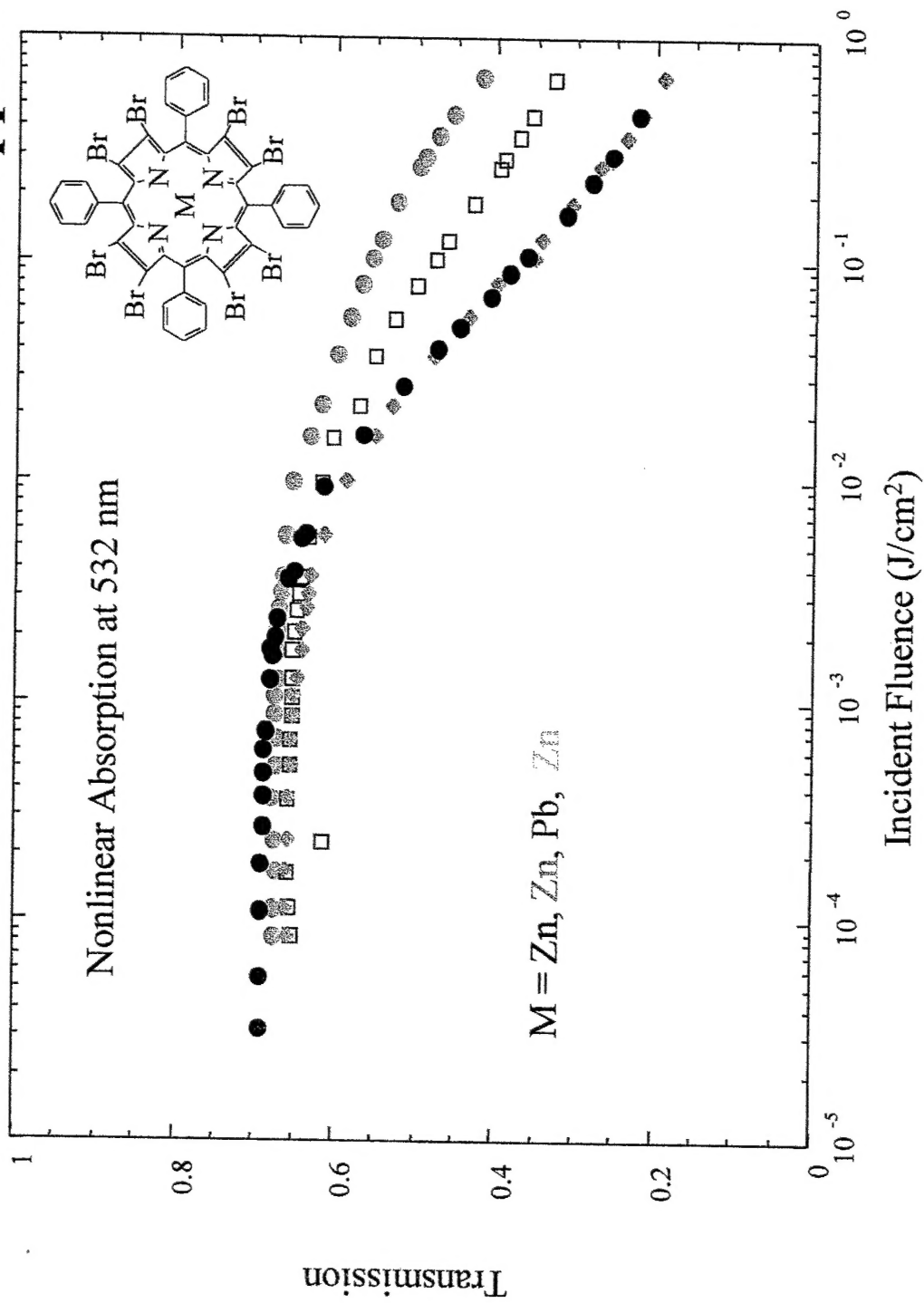
- determine barriers to polymerization of ethylene, oxidation by HO_2H .



PROJECT OVERVIEW - NLO



Non-linear optical materials for laser-hardened applications



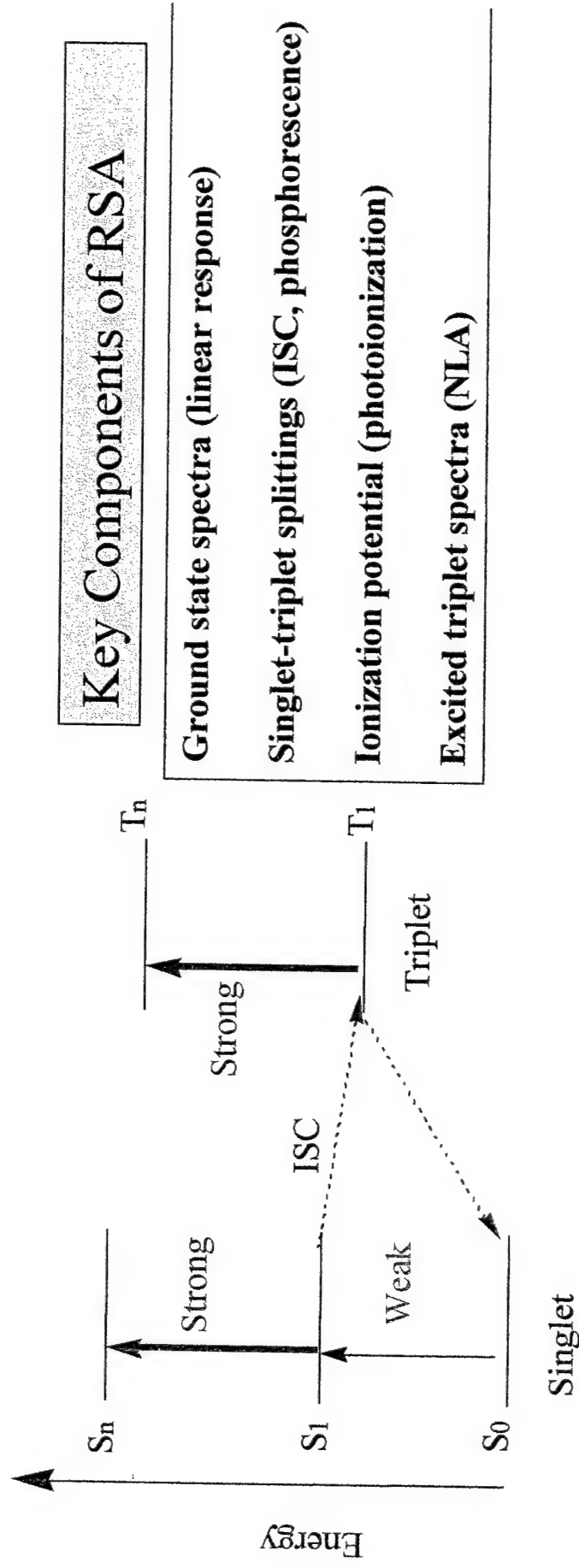


PROJECT OVERVIEW - NLO

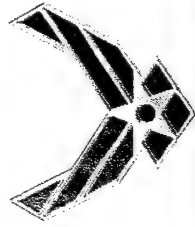


Technical issues being addressed using CCM

1. Mechanism of reverse saturable absorption (RSA)



Five-level model for nonlinear absorption



THEORETICAL METHODS

1. Ab initio electronic structure theory

- General Atomic and Molecular Electronic Structure System (GAMESS) -- a CHSSI code
- Nuclear-electronic orbital approach (NEO) for including nuclear quantum effects (important, e.g., in proton transfer reactions)

Various computational techniques are employed to solve the molecular electronic Schrödinger equation from quantum mechanics:

$$\left[-\frac{1}{2} \sum_i \nabla_i^2 - \sum_i \sum_{\alpha} \frac{Z_{\alpha}}{r_{i\alpha}} + \sum_i \sum_{j>i} \frac{1}{r_{ij}} \right] \Psi_{el} = E_{el} \Psi_{el}$$

Categories of approximate solutions:

- a) “Self-consistent field” (SCF): reasonably good geometries
- b) “Electron correlation”: post-SCF correction, required for reliable energetics (e.g., barriers).



THEORETICAL METHODS

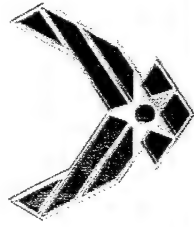
1. Ab initio electronic structure theory (cont.)

- Most electronic structure codes use Born-Oppenheimer (i.e., “clamped nuclei”) approximation -- NOE method treats specified nuclei at QM level.

Nuclear-Electronic Hamiltonian

$$H_{\text{tot}}(\mathbf{r}_e, \mathbf{r}_q; \mathbf{r}_c) = -\sum_i \frac{N_e}{2} \nabla_i^2 - \sum_i \sum_A \frac{N_e N_c Z_A}{r_{iA}} + \sum_i \sum_{j>i} \frac{N_e N_e}{r_{ij}} \\ - \sum_I \frac{N_p}{2M_I} \nabla_I^2 + \sum_I \sum_A \frac{N_p N_c Z_A Z_I}{r_{IA}} + \sum_I \sum_{J>I} \frac{N_p N_p Z_I Z_J}{r_{IJ}} \\ - \sum_i \sum_I \frac{N_e N_p Z_I}{r_{iI}} + \sum_A \sum_{B>A} \frac{N_c N_c Z_A Z_B}{r_{AB}}$$

- N_e : number of electrons (coordinates \mathbf{r}_e)
 N_p : number of quantum nuclei (coordinates \mathbf{r}_p)
 N_c : number of classical nuclei (coordinates \mathbf{r}_c)



THEORETICAL METHODS



Ab initio electronic structure theory

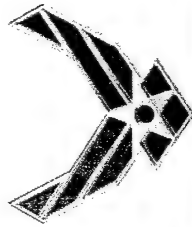
Current Status of parallel GAMESS

	<u>RHF</u>	<u>ROHF</u>	<u>UHF</u>	<u>GVB</u>	<u>MCSCF</u>
Energy	cdp	cdp	cdp	cdp	cdp
Analytic Gradient	cdp	cdp	cdp	cdp	cdp
Numeric Hessian	cdp	cdp	cdp	cdp	cdp
Analytic Hessian	cdp	cdp	-	cdp	-
MP2 energy	cdp	cdp	cdp	-	c p
MP2 gradient	cdp	-	cd	-	-
CI energy	cdp	cdp	-	cdp	cdp
CI gradient	cd	-	-	-	-
DFT energy	cdp	cdp	cdp	-	-
DFT gradient	cdp	cdp	cdp	-	-

c = conventional disk storage of AO integrals

d = direct evaluation of AO integrals

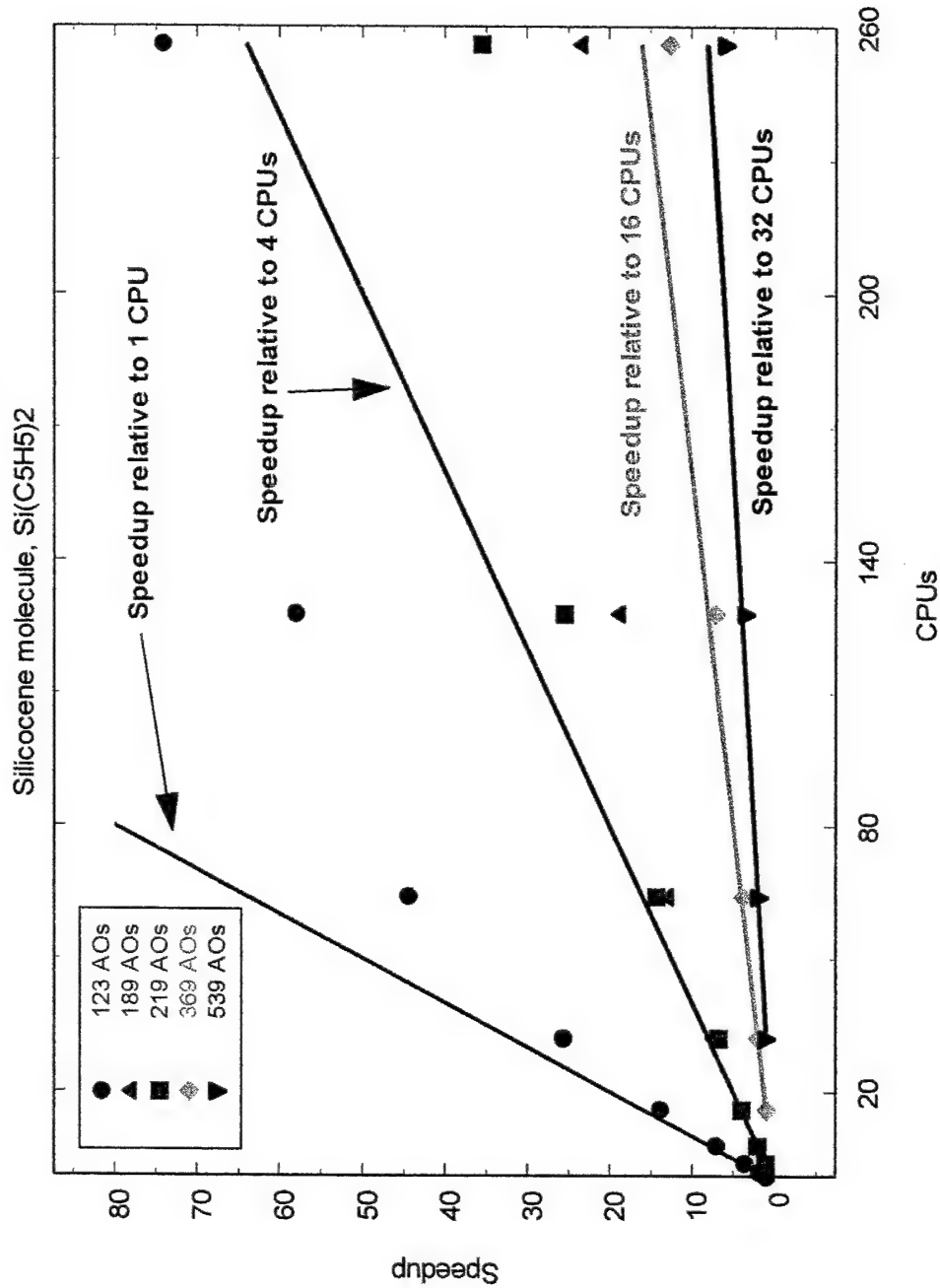
p = runs in parallel

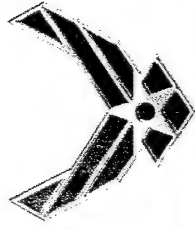


THEORETICAL METHODS

Ab initio electronic structure theory

MP2 Gradient Scalability Test

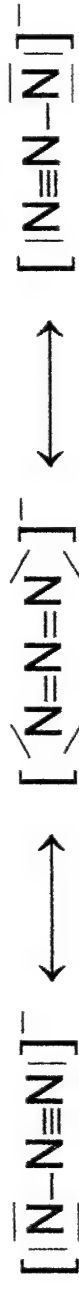




The Search for New Polynitrogens



- All polynitrogens are unstable with respect to N_2 molecules
- Their activation energy for N_2 elimination is largely determined by the weakest bond in the compound
- Their metastability is enhanced if suitable resonance structures exist:



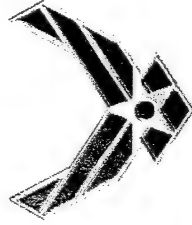
- The double-bond character of the N—N bonds in the azide anion explains its exceptional stability
- How can this stabilization effect be used to our advantage in preparing new compounds?



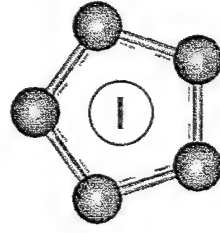
Pentazolate (N_5^-)?



- Substituted pentazoles $R-N_5$ have been known for decades ($R=aryl$)
- Cyclic N_5^- is aromatic
- Conversion of the diazonium salt, RN_2^+ , to the substituted pentazole ring $R-N_5$ by the reaction with azide ion, N_3^- , has been demonstrated many years ago by Ugi and Huisgen.
- N_5^- has been recently detected in the gas phase for the first time, using collisional fragmentation (electrospray ion mass spectroscopy).
- Can a chemical route to N_5^- be found? (e.g., can a suitable R group be found for the reaction $R-N_2^+ + N_3^- \rightarrow R-N_5 \rightarrow R^+ + N_5^-$?)

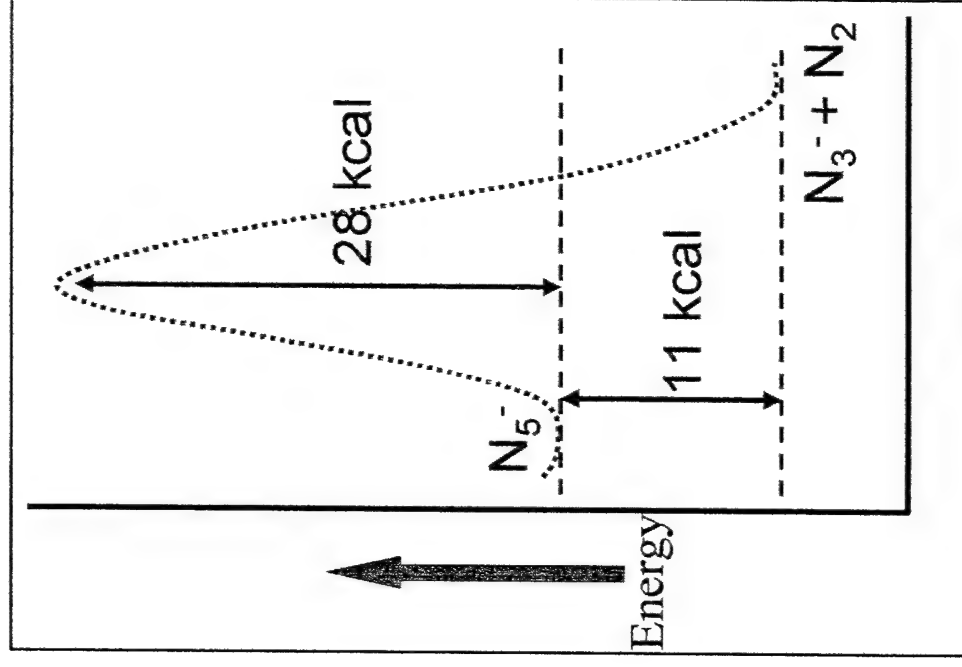


New Polynitrogen Anions



Pentazole anion (N_5^-)

- Theoretical calculations show that this anion has a 28 kcal/mole activation energy barrier for decomposition and its decomposition to N_3^- and N_2 is only 11 kcal/mol exothermic
- Aryl substituted pentazoles can be isolated as stable compounds only if stored at low temperatures. In methanol, these compounds rapidly decompose at room temperature to form aryl azides and N_2 gas





Synthetic Challenge – How do we make These New Anions??



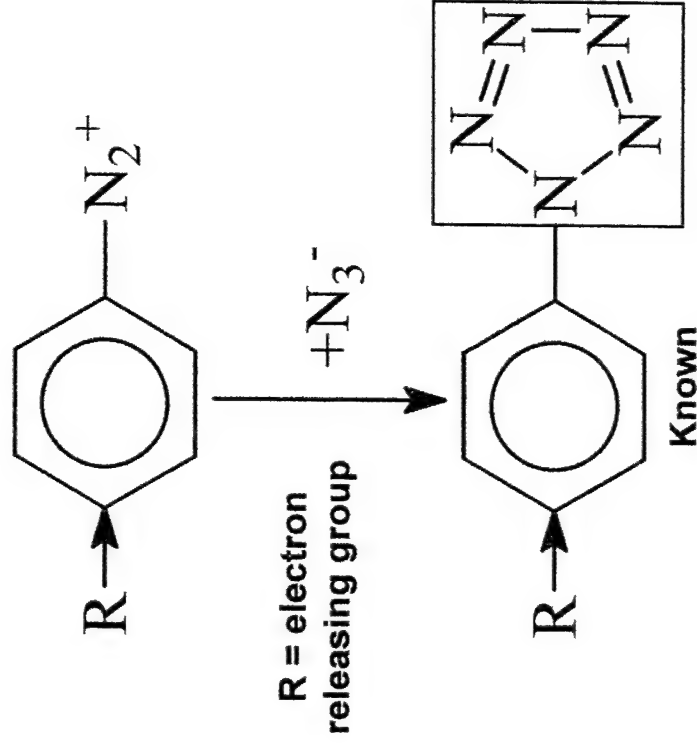
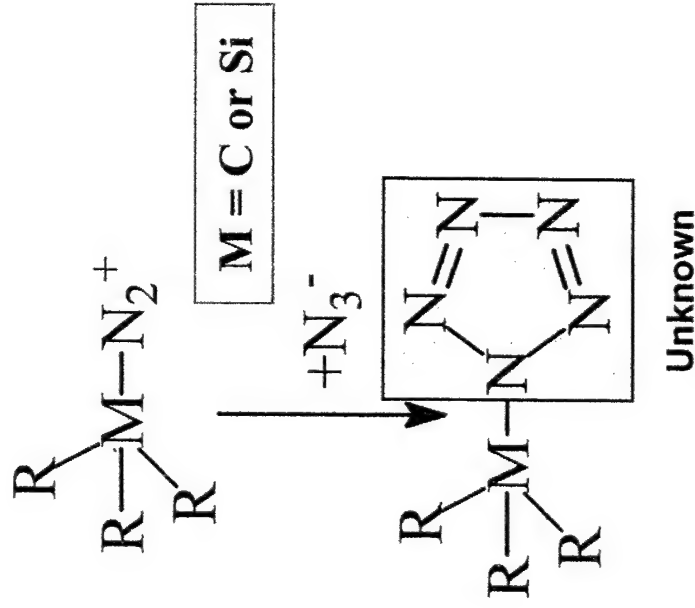
Synthesis of Substituted Pentazoles

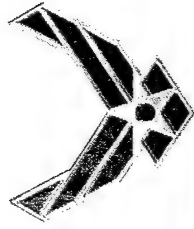
Sources for the Pentazole Anion (N_5^-)



Trityl/Silyl Diazonium Salts

Aryl Diazonium Salts





Theoretical Challenge - Can we design and predict viable precursors to N_5^- ?



Find a substituent R so that

1. $R-N_2^+$ is stable wrt $R^+ + N_2$.
2. The R-N bond in $R-N_5$ is weak, thereby suitable for the reaction $R-N_5 + M^+X^- \rightarrow M^+N_5^- + R-X$

Approach

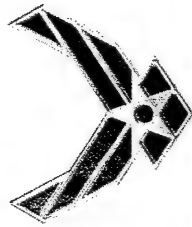
Use quantum chemical calculations to predict the stability of $R-N_2^+$ diazonium salts and the length/strength of the $R-N_5$ bond.

MBPT(2)/6-31G(d) level of theory



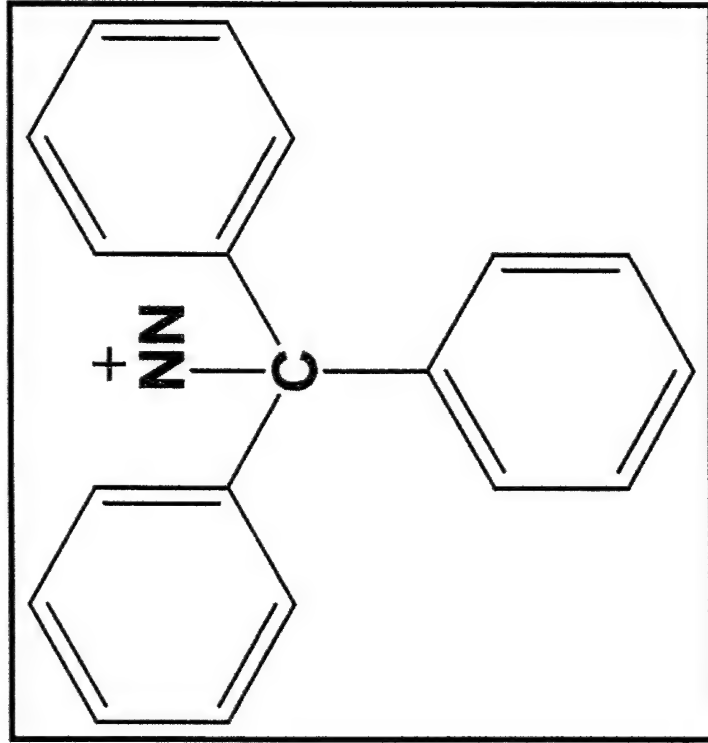
M = C, Si

L = F, Cl, CH_3 , CF_3 , NO_2 , phenyl, etc.

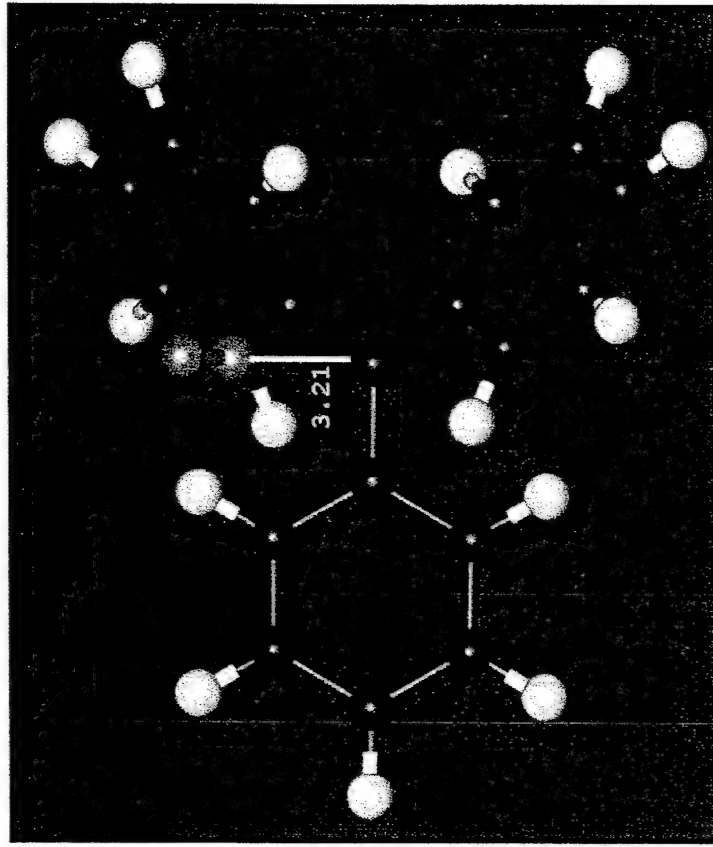


Identifying Precursors for New Polynitrogens

*This ion has been suggested
as a useful precursor to new
polynitrogen molecules...*



*... but calculations predict it to be
unstable.*



Computational requirements: ~50,000 CPU-hours, 1.2 GW on IBM SP/P3 at ASC

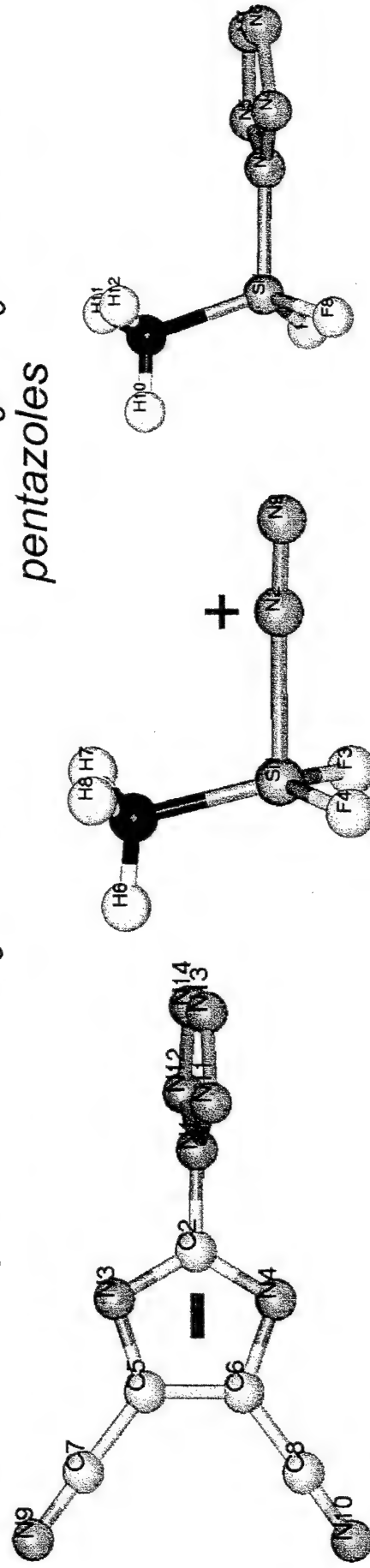


RESULTS - HEDM



High-nitrogen/polynitrogen compounds

Potential dicyanoimidazolate $[R_3Si-N_2]^+$ precursors to precursor to displacement of N_5^- formation of suitable R_3Si-N_5 substituted pentazoles



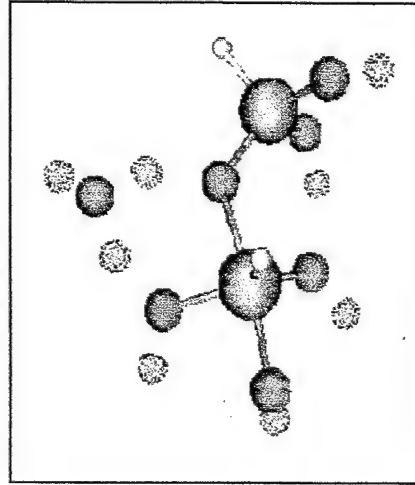
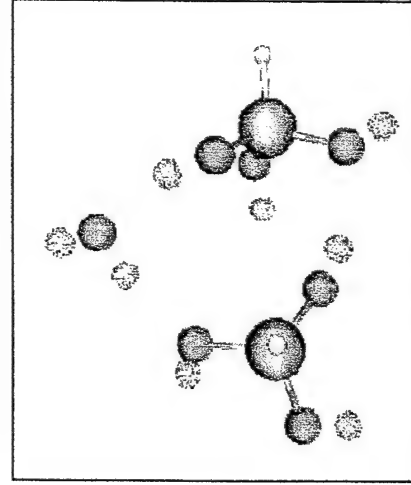
R_3	$R_3Si-N_2^+$ distance(Å)	$D_e(Si-N_2^+)$ (kcal/mol)	R_3Si-N_5 distance (Å)
3Me	2.151	13.8	1.855
2Me,F	2.144	15.7	1.834
Me,2F	2.151	21.2	1.855
3F	1.973	33.4	1.783

Computational requirements: ~50,000 CPU-hours, 1 GW on IBM SP/P3 at ASC

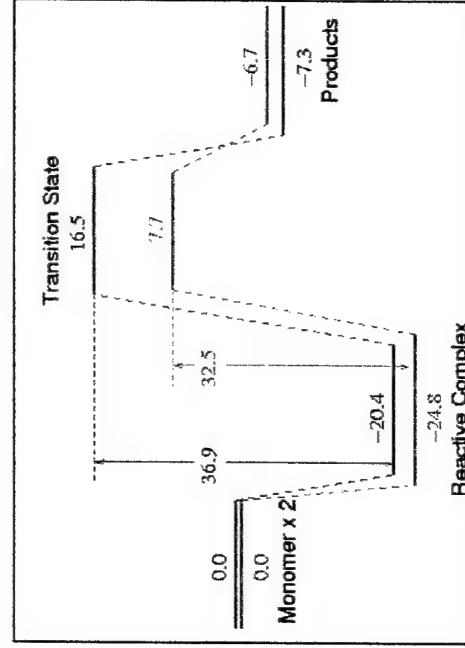
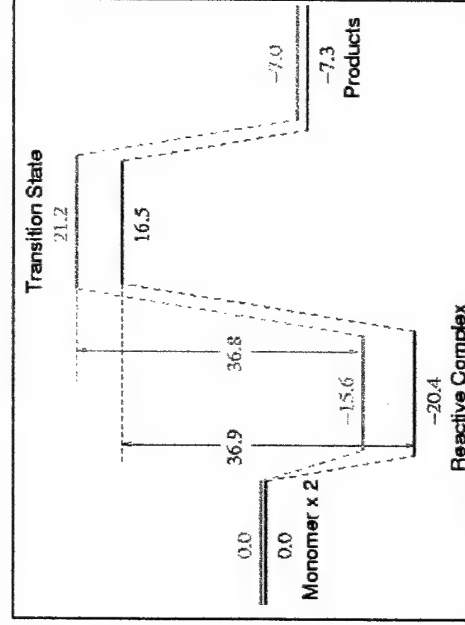


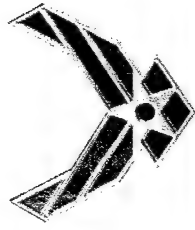
RESULTS - POSS

Nuclear quantum effects in water-catalyzed condensation reactions



Level	Reactive Complex	Transition State	Products	CPU Time
RHF/6-31G*	-20.4	16.5	-7.3	1.0
RHF/6-31G*/ZPE	-15.6	21.2	-7.0	
NEO/HF/4	-23.0	7.2	-5.9	1.1
NEO/HF/8	-24.8	7.7	-6.7	1.8





POSS as catalysts?

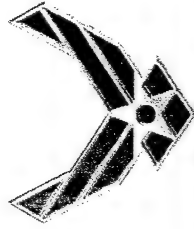


Some Ti compounds are known to be effective catalysts

Ziegler-Natta, olefin oxidation

Recent experiments reveal Ti-modified silicates catalyze olefin oxidation by peroxides

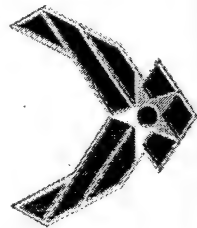
Can Ti-substituted siloxanes and/or POSS catalyze olefin oxidation by HOOH , or polymerization?



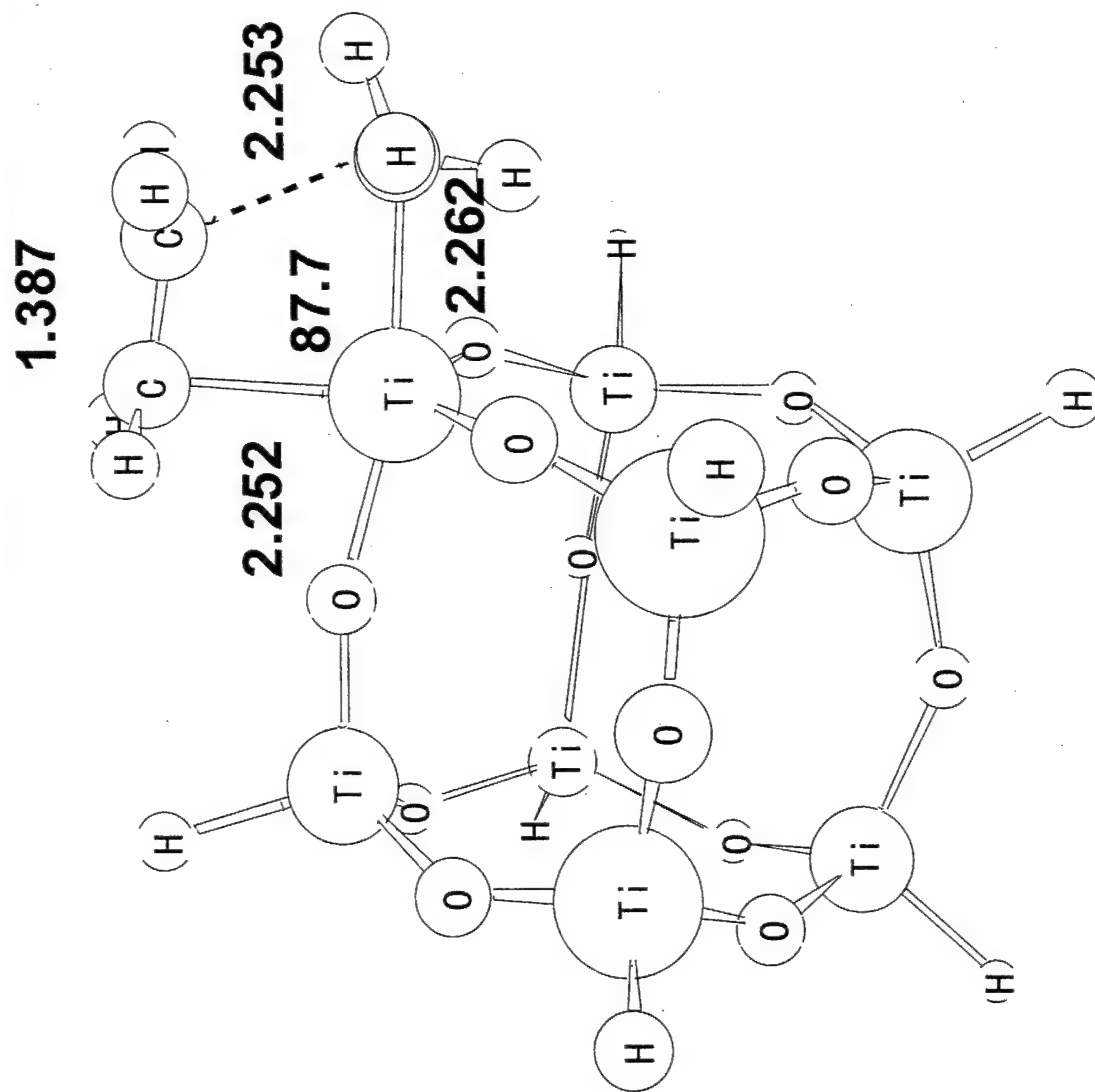
POSS computational methods

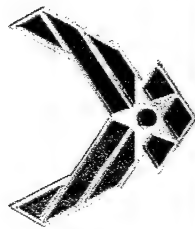


- § B3LYP/6-31G(d) geometry optimizations
- § Hessians to characterize stationary points
- § Energies using MP2/TZVP
- § Some geometries re-optimized with MP2
- § Size of basis set: ~1,000 basis functions
- § Calculations performed at ERDC (T3E) and AHPCRC (T3E)



TS For Polymerization of Ethylene





RESULTS - NLO



Systematic investigation of excited state energies and oscillator strengths of free-base porphyrins, phthalocyanines, and their metal complexes, representing a broad range of π -conjugated systems, using time-dependent density functional theory (TD-DFT).

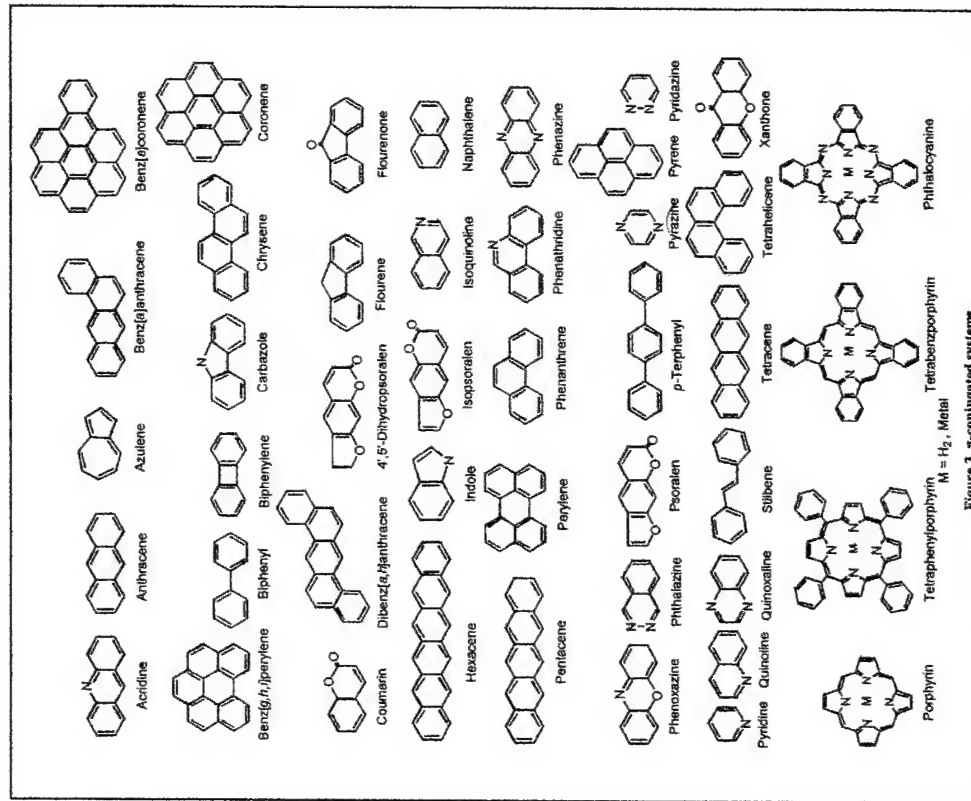
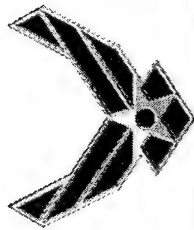
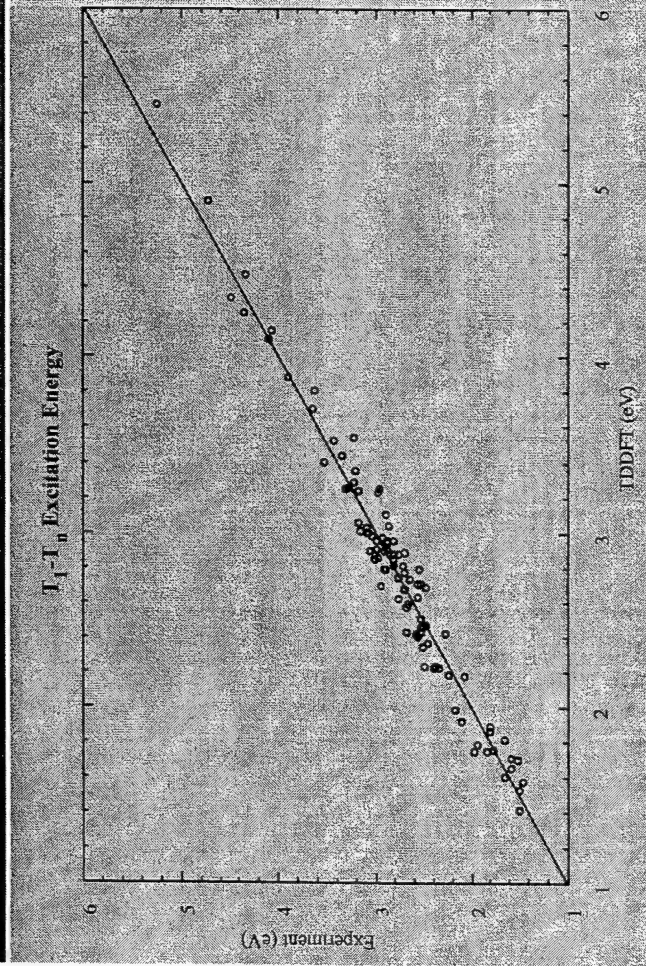


Figure 3. π -conjugated systems

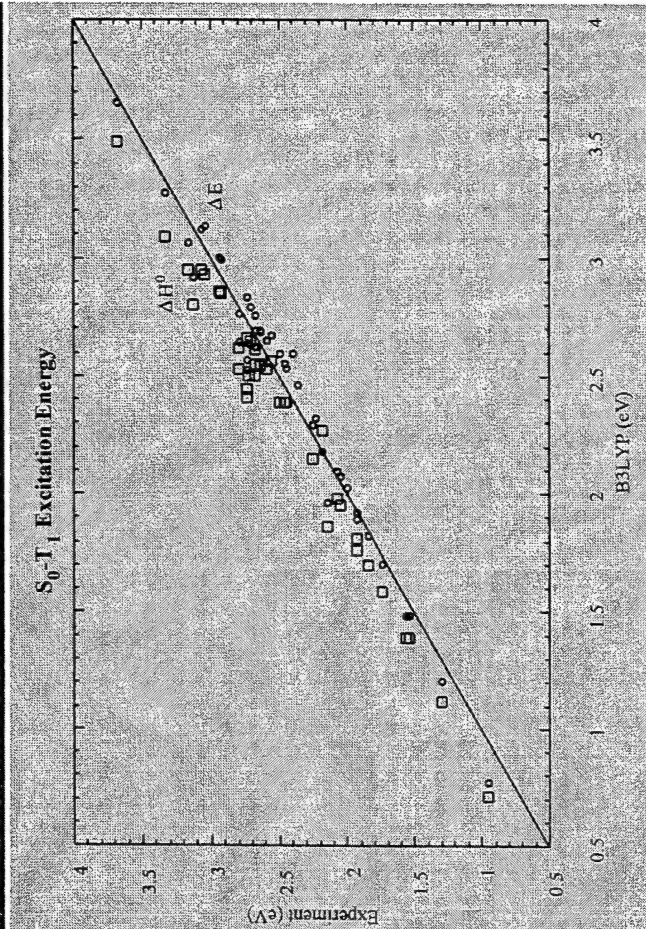


RESULTS - NLO



Computed T_1-T_n compared to experimental results

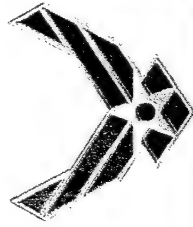
86 Experimental Values (in solution)
Mean Absolute Error = 0.11
Maximum Error = 0.31



Computed S_0-T_1 compared to experimental results

47 Experimental Values (in solution)
Mean Absolute Error = 0.14
Maximum Error = 0.33

100,000 hours on SGI O2K, Compaq GS320, IBM SP/P3 at ASC



SUMMARY

High Energy Density Matter

- High-nitrogen/polynitrogen compounds are substantially more energetic than hydrazine.
- Trityldiazonium cation is not a stable polynitrogen precursor.
- Trimethylsilyldiazonium cation is marginally stable, replacement of methyls with fluorines increases stability.

Polyhedral Oligomeric Silsesquioxanes (POSS)

- Nuclear quantum effects are important in proton transfer reactions -- lower barriers by about 4 kcal/mol for water-assisted condensation of trisilanol
- Most effective catalyst for olefin polymerization is T_4 cage
- Barrier is still ~10 kcal/mol too high: Ti-POSS, Ti-siloxanes may not be good catalysts for this reaction

NLO materials

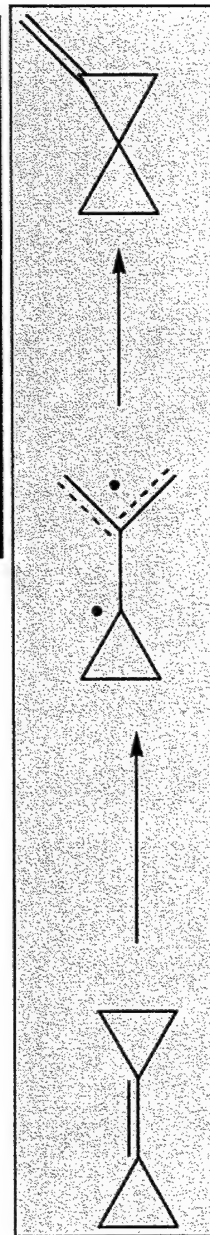
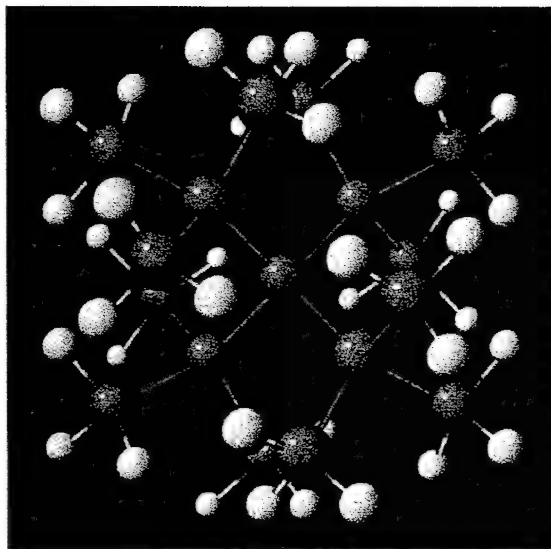
- Time-dependent density functional theory accurately predicts NLA in free-base porphyrins, phthalocyanines, and their metal complexes.
- Mean absolute error of 0.11 eV for computed triplet-triplet excitation energies
- Mean absolute error of 0.14 eV for computed singlet-triplet excitation energies

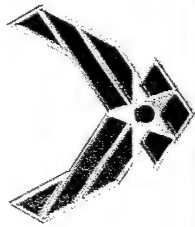


FUTURE DIRECTIONS

HEDM

- Thermal decomposition mechanisms of energetic hydrocarbons (e.g., BCP).
- Energetic Ionic Liquids (QSPR models)





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GAMESS: Graham Fletcher

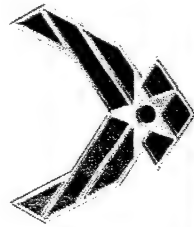
MSRCs, DCs: ASC, ARL, ERDC, NAVO, MHPCC, AHPCRC, ARSC, AFFTC

CHSSI funding (CCM-2, CCM-4, MBD-01)



Backup Slides





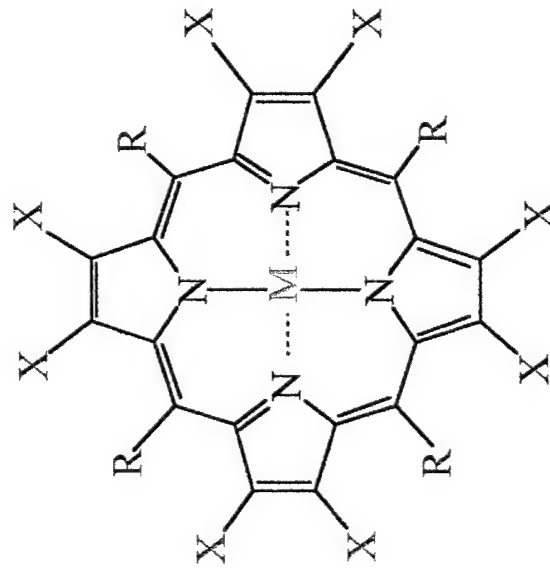
PROJECT OVERVIEW - NLO



Technical issues being addressed using CCM

2. "Tuning" of absorption spectrum by benzannulation, halide substitution

System	Property						
	M	X	R	IP	S ₀ -S _n	S ₀ -T ₁	T ₁ -T _n
PH ₂	H ₂	H	H	E, C	E, C	E, C	E, C
ZnP	Zn	H	H	E, C	E, C	E, C	E, C
TPPH ₂	Zn	H	φ	E, C	E, C	E, C	E, C
ZnTPP	Zn	H	φ	E, C	E, C	E, C	E, C
ZnTPPBr ₈	Zn	Br	φ	C	E, C	E, C	E, C



IP = Ionization Potential, S₀-S_n = Ground State Spectrum,
 S₀-T₁ = Singlet-Triplet Gap T₁-T_n = Triplet-Triplet Spectrum
 E = Experiment, C = Calculated



RESULTS - POSS

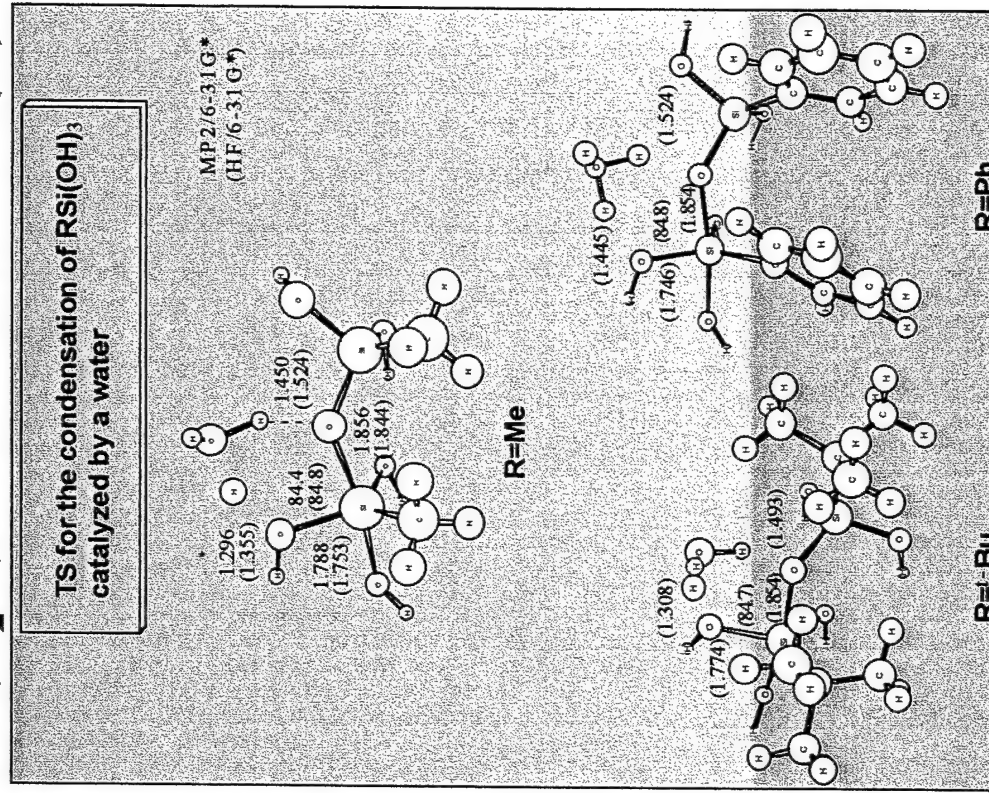


Mechanism of formation: role of solvent (H_2O) & substituents (R)

$\text{RSi}(\text{OH})_3 + \text{RSi}(\text{OH})_2 + \text{H}_2\text{O} \rightarrow \text{RSi}(\text{OH})_2\text{Si}(\text{OH})_2 + \text{H}_2\text{O}$			
R	Energy barrier (kcal/mol)		MP2/6-31G*
	HF/6-31G*	MP2/6-31G*	
H	30.4 (16.7)	10.9 (-9.3)	
Me	28.2 (14.7)	7.7 (-13.3)	
t-Bu	34.3 (24.9)	9.8 (-9.3)	
Ph	31.1 (18.2)	7.9 (-16.4)	

Values in parentheses are for water-catalyzed results.

Kudo, T., Gordon, M.S. J. Am. Chem. Soc., 120, 11432 (1998)
Kudo, T., Gordon, M.S. J. Phys. Chem. A, 104, 4058 (2000)





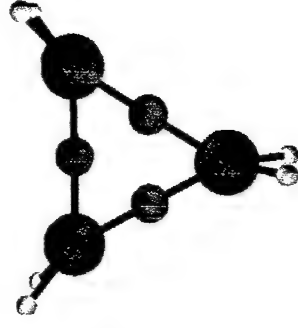
RESULTS - POSS



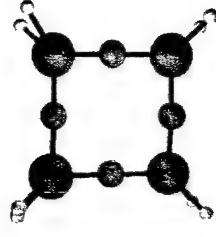
Mechanism of formation

Key steps

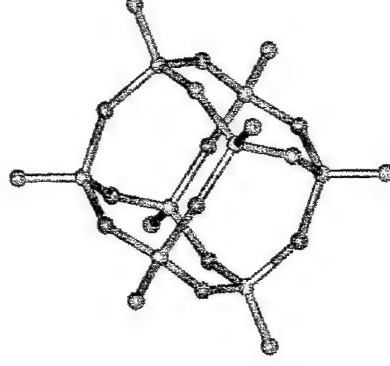
1. Hydrolysis of RSiX_3 ($\text{R}=\text{H}, \text{CH}_3, \text{t-butyl}, \text{etc.}; \text{X}=\text{Cl}$)
 $\text{RSiCl}_3 + \text{H}_2\text{O} \rightarrow \text{RSiCl}_2\text{OH} + \text{HCl}$
 $\text{RSiCl}_2\text{OH} + \text{H}_2\text{O} \rightarrow \text{RSiCl(OH)}_2 + \text{HCl}$
 $\text{RSiCl(OH)}_2 + \text{H}_2\text{O} \rightarrow \text{RSi(OH)}_3$
2. Condensation of RSi(OH)_3 to disiloxane
 $2 \text{RSi(OH)}_3 \rightarrow \text{R(OH)}_2\text{SiOSi(OH)}_2\text{R} + \text{H}_2\text{O}$
3. Condensation of disiloxane to D_3, D_4
 $\text{RSi(OH)}_3 + \text{R(OH)}_2\text{SiOSi(OH)}_2\text{R} \rightarrow \text{D}_3 + 2\text{H}_2\text{O}$
 $[2+2]: 2\text{R(OH)}_2\text{SiOSi(OH)}_2\text{R} \rightarrow \text{D}_4 + 2\text{H}_2\text{O}$
 $[3+1]: \text{RSi(OH)}_3 + \text{R(OH)}_2\text{SiOSi(OH)}_2\text{R} \rightarrow \text{D}_4 + 2\text{H}_2\text{O}$
Ring Expansion: $\text{RSi(OH)}_3 + \text{D}_3 \rightarrow \text{D}_4 + \text{H}_2\text{O}$
4. Condensation of D_3, D_4 to POSS (in progress)
 $2\text{D}_4 \rightarrow \text{T}_8 + 4\text{H}_2\text{O}$
....



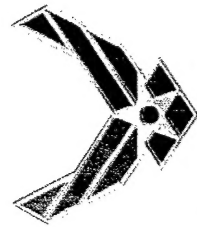
D3



D4

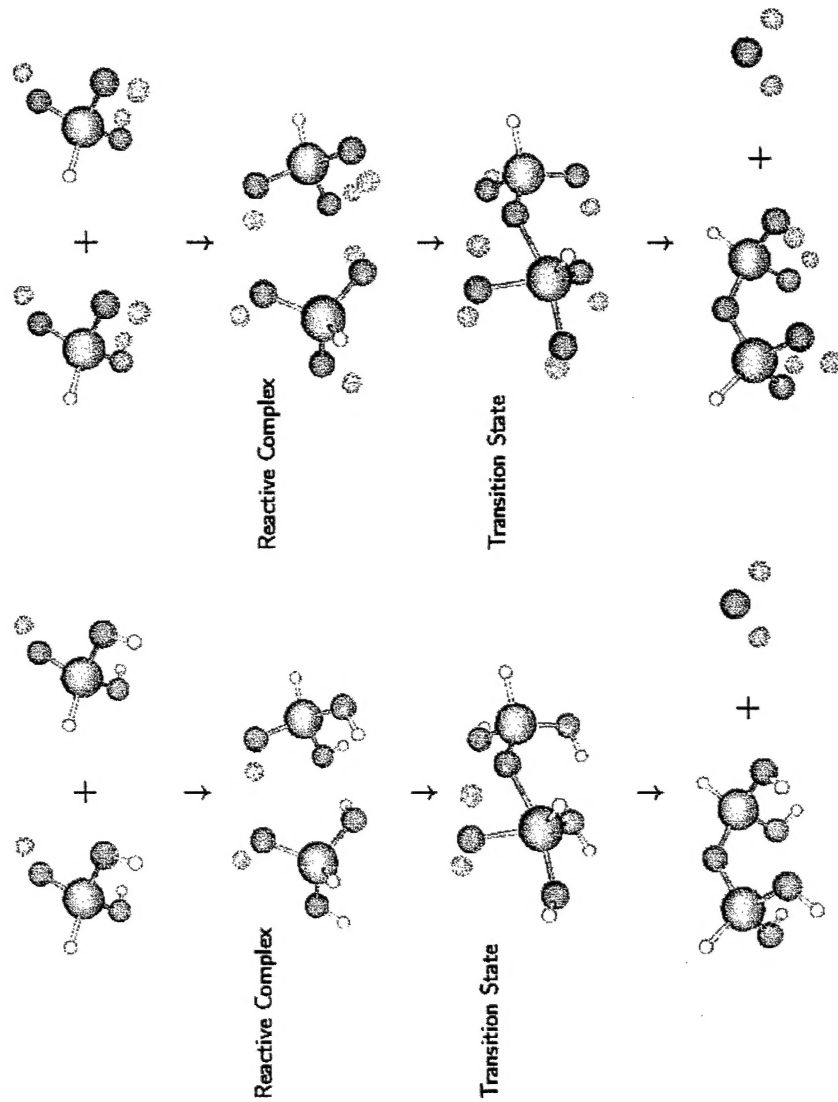


T8



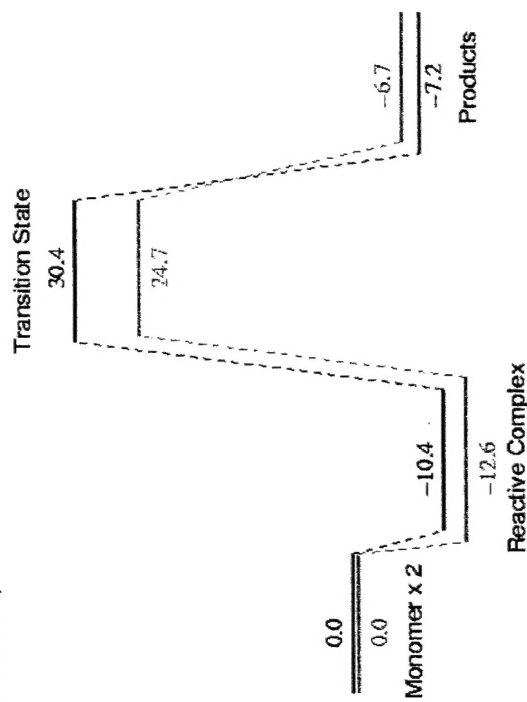
RESULTS - POSS

Nuclear quantum effects in condensation reactions



Level	Monomer $\times 2$	Reactive Complex	Transition State	Products	CPU Time
RHF/6-31G*	0.0	-10.4	30.4	-7.2	1.0
NEO-HF/2	0.0	-11.4	24.9	-5.9	1.1
NEO-HF/6	0.0	-12.6	24.7	-6.7	1.8

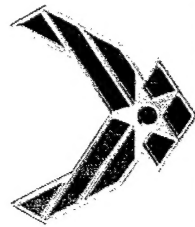
Units in kcal/mol



Hammes-Schiffer, S., J. Phys. Chem. A 102 (1998), 10443

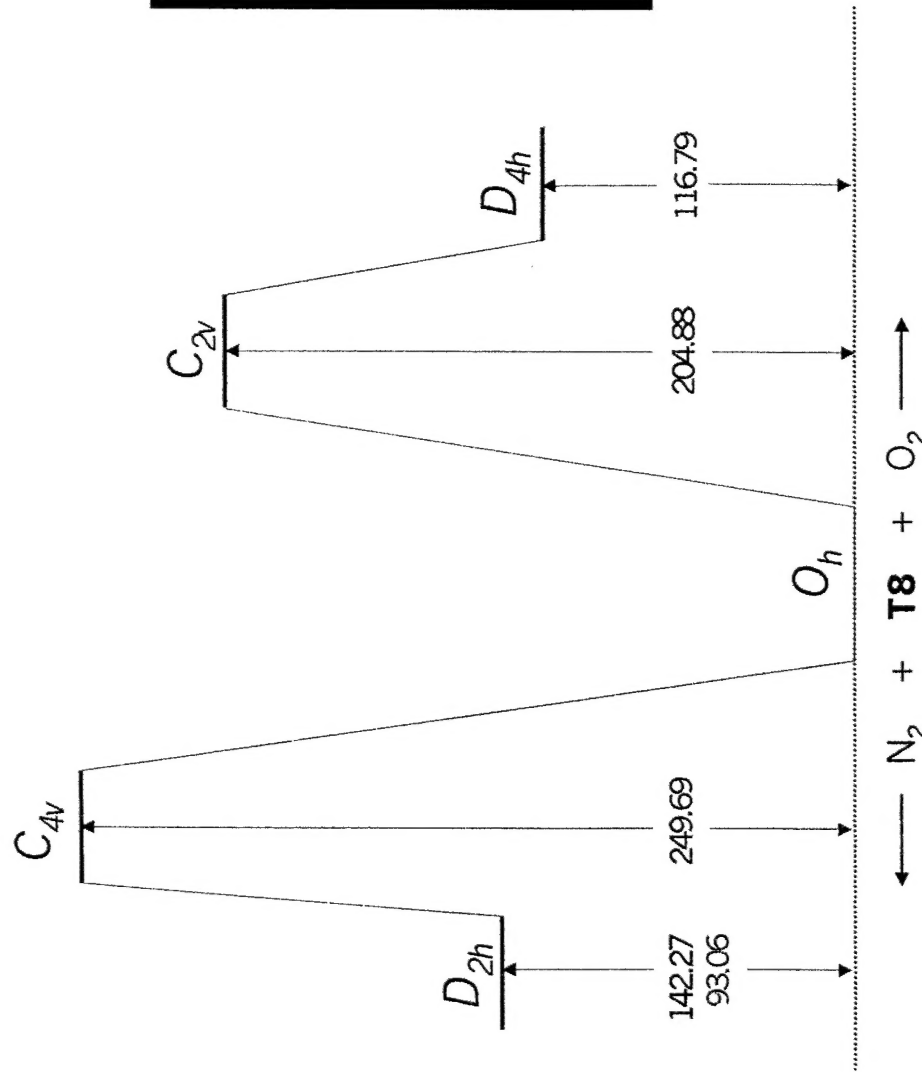
Webb, S.P., Agarwal, P.K., and Hammes-Schiffer, S., J. Phys. Chem. B, 104(2000), 888

Webb, S.P. and Hammes-Schiffer, S., J. Chem. Phys. 113 (2000), 5214

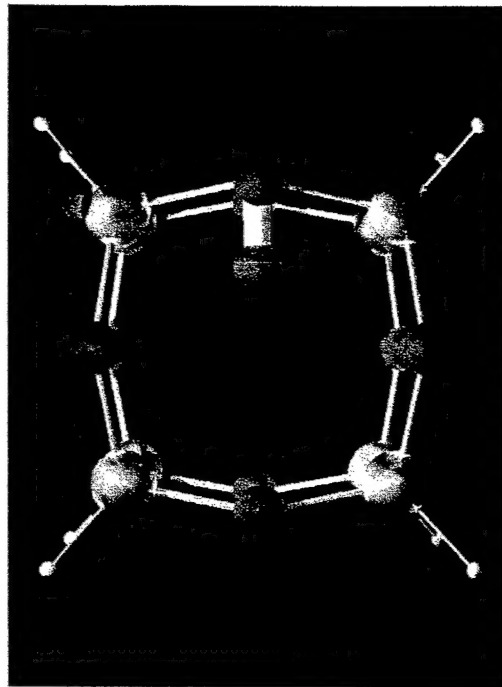


RESULTS - POSS

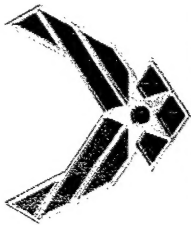
Molecular "sieves": preferential capture N_2 vs. O_2 ?



TS structure of $T_8 + O_2$



T_{10} and T_{12} calculations in progress
HPC requirements: ~50,000 node-hrs, AHPARC T3E, 256 GB



RESULTS - NLO



B3LYP S_0-T_1 Excitation Energies (in eV)

System	6-31G(d)	Error	Exp
Porphyrin (1^3B_{2u})	1.42	0.16	1.58 ^a
Zinc Porphyrin (1^3B_{1u})	1.65	0.07	1.72 ^b
Tetraphenylporphyrin (1^3B_1)	1.31	0.14	1.45 ^c
Zinc Tetraphenylporphyrin(1^3B_1)	1.53	0.06	1.59 ^d
Zinc Phthalocyanine (1^3B_{2u})	1.05	0.08	1.13 ^e
Zinc Tetrabenzporphyrin (1^3B_{1u})	1.41	0.16	1.57 ^f
Phthalocyanine (1^3B_{1u})	1.18	0.06	1.24 ^g
Mean Error		0.10	

^aGouterman, Khalil, *J. Mol. Spectrosc.* 1974, 53, 88. (EPA (5:5:2) mixture of ethyl ether to isopentane to ethanol) and 50% ethyl iodide at 77 K) ^bGradyushko, Tsvirko, *Opt. Spectrosc.* 1971, 31, 291.(EPA at 77 K) ^cGouterman, Khalil, *J. Mol. Spectrosc.* 1974, 53, 88. (EPA at 77 K) ^dWalters et al., *J. Phys. Chem.* 1995, 99, 1166.(1:1 mixture of ether to ethanol at 77 K) ^eVincett et al.,K. E. *J. Chem. Phys.* 1971, 55, 4131. (1-chloronaphthalene at 77 K) ^fBajema, Gouterman, *J. Mol. Spectrosc.* 1971, 39, 421 (octane at 77 K) ^gMcVie et al., *J. Chem. Soc. Faraday Trans. II* 1978, 74, 1870 (1-chloronaphthalene at 77 K)

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Nguyen, K. A., Day, P. N., and Pachter, R., *J. Phys. Chem. A*, 103 (1999) 7378

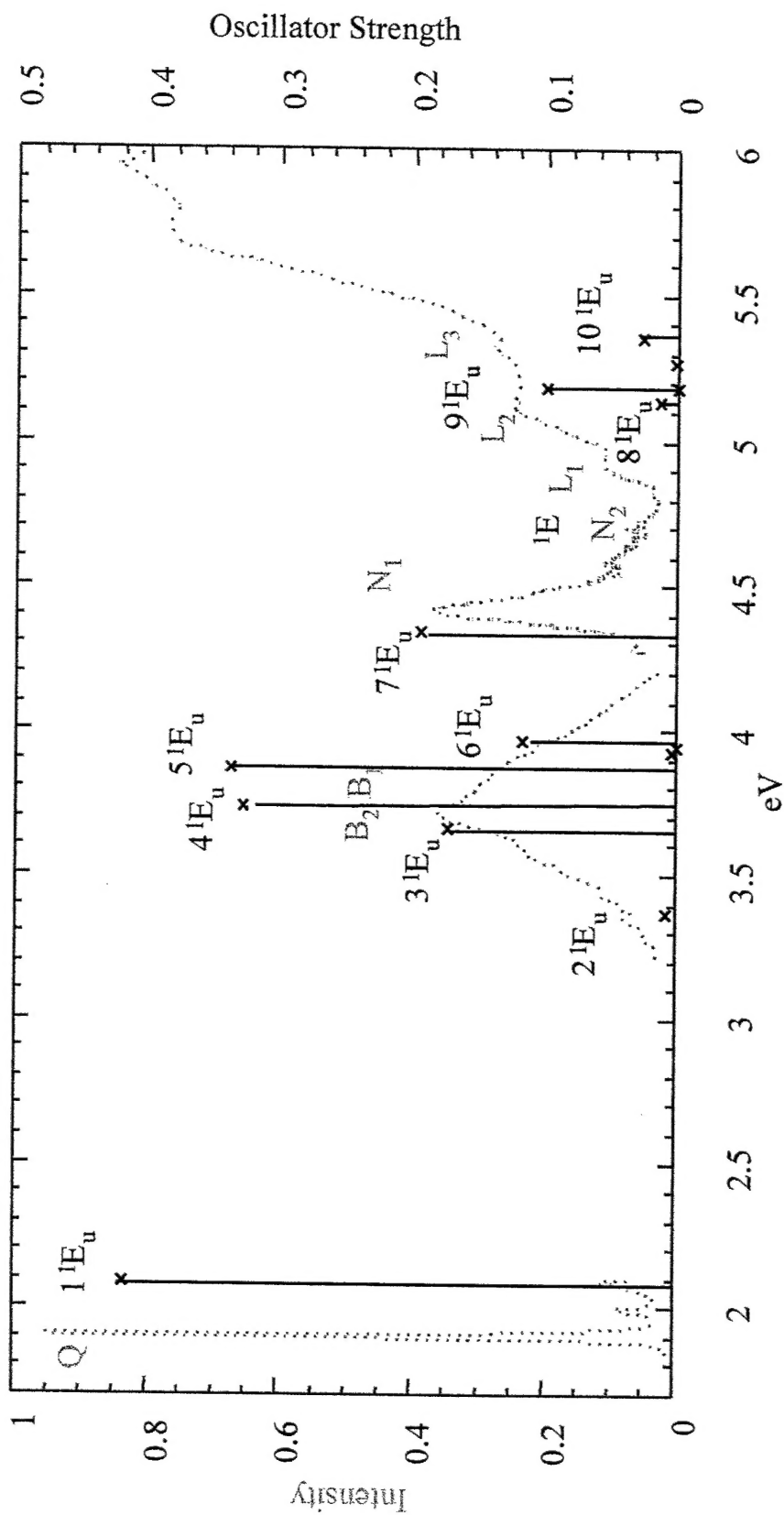
Nguyen, K. A., Day, P. N., and Pachter, R., *J. Phys. Chem.*, 104 (2000) 4755

Nguyen, K. A. and Pachter, R., *J. Phys. Chem.*, 104 (2000) 4549



RESULTS - NLO

Comparison with Experiment: ZnPc



HPC Requirements: ~100,000 CPU hours, on SGI O2K + IBM SP3 + SGI O2K @ ASC